=> Uploading 09319967.str

STRUCTURE UPLOADED L6

=> d

L6 HAS NO ANSWERS

Ь6

0

Structure attributes must be viewed using STN Express query preparation.

=> s 2/si

112281 2/SI L7

=> s 16 subset=17

ENTER SUBSET SEARCH SCOPE - SAMPLE, FULL, RANGE, OR (END):sam

SAMPLE SUBSET SEARCH INITIATED 10:29:01 FILE 'REGISTRY' SAMPLE SUBSET SCREEN SEARCH COMPLETED - 2 TO ITERATE

100.0% PROCESSED 2 ITERATIONS 1 ANSWERS

SEARCH TIME: 00.00.01

PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE \*\*COMPLETE\*\*

PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 2 TO 124

1 TO PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET): 80

1 SEA SUB=L7 SSS SAM L6 L8

=> s scan

Ь9 14 SCAN

=> d scan 18

REGISTRY COPYRIGHT 2000 ACS L8 1 ANSWERS

Tyrosine, 3,5-dimethoxy-O-methyl-N-(trifluoroacetyl)-N-(trimethylsilyl)-IN

 $\beta$ -[(trimethylsilyl)oxy]-, methyl ester, erythro- (9CI)

C21 H34 F3 N O7 Si2 MF

Relative stereochemistry.

## ALL ANSWERS HAVE BEEN SCANNED

=> s 16 subset=17 full

FULL SUBSET SEARCH INITIATED 10:29:48 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 79 TO ITERATE 100.0% PROCESSED 79 ITERATIONS

75 ANSWERS

SEARCH TIME: 00.00.02

75 SEA SUB=L7 SSS FUL L6

=> s 110 and 1/o

2607474 1/0

19 L10 AND 1/0 L11

=> d scan

L11 19 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN L-Proline, 1-L-alanyl-, mixt. with trimethylsilyl 2,2,2-trifluoro-N-(trimethylsilyl)ethanimidate (9CI)

C8 H18 F3 N O Si2 . C8 H14 N2 O3 MF

CI MXS

> CM1

$$\begin{array}{c} \text{O-SiMe}_3\\ |\\ \text{Me}_3\text{Si-N----} \text{C-CF}_3 \end{array}$$

CM2

Absolute stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

 $\Rightarrow$  s 111 and 1/nc

19101843 1/NC L12 14 L11 AND 1/NC

=> d scan

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Ethanimidic acid, N-(dimethylpropylsily1)-2,2,2-trifluoro-,
dimethylpropylsily1 ester (9CI)
MF C12 H26 F3 N O Si2

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):5

L12 14 ANSWERS REGISTRY. COPYRIGHT 2000 ACS
IN Silanamine, 1,1,1-trimethyl-N-[2,2,2-trifluoro-1[(trimethylsilyl)oxy]ethyl]- (9CI)
MF C8 H20 F3 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Silanamine, N-[1-[(dimethyl-2-propenylsilyl)oxy]-2,2,2-trifluoroethyl]-1,1dimethyl-1-(2-propenyl)- (9CI)
MF C12 H24 F3 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, 2,2,2-trifluoro-N-(triethylsilyl)-, triethylsilyl ester (9CI)

MF C14 H30 F3 N O Si2

$$\begin{array}{c} \text{O-SiEt}_3 \\ | \\ \text{Et}_3 \text{Si-N---} \text{C-CF}_3 \end{array}$$

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Silanamine, 1,1,1-trimethyl-N-[2,2,2-trifluoro-1-(trifluoromethyl)-1-[(trimethylsilyl)oxy]ethyl]- (9CI)

MF C9 H19 F6 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS

MF C14 H30 F3 N O Si2

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):10

14 ANSWERS REGISTRY COPYRIGHT 2000 ACS L12

Ethanimidic acid, N-(diethylsilyl)-2,2,2-trifluoro-, diethylsilyl ester

MF C10 H22 F3 N O Si2

$$\begin{array}{c} \text{O-SiHEt}_2 \\ | \\ \text{Et}_2 \text{SiH-N---} \text{C-CF}_3 \end{array}$$

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS Ethanimidic acid, N-(dimethyl-2-propenylsilyl)-2,2,2-trifluoro-, dimethyl-2-propenylsilyl ester (9CI)

C12 H22 F3 N O Si2 MF

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS

Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl) - (8CI, 9CI) IN

C8 H18 F3 N O Si2 MF

CI COM

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS

Ethanimidic acid, N-(ethyldimethylsilyl)-2,2,2-trifluoro-,

ethyldimethylsilyl ester (9CI)

MF C10 H22 F3 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Acetamide, N,N-bis(dimethyl-2-propenylsilyl)-2,2,2-trifluoro- (9CI)
MF C12 H22 F3 N O Si2

$$\begin{array}{c} \text{Me} \\ | \\ \text{H}_2\text{C} \longrightarrow \text{CH} - \text{CH}_2 - \text{Si} - \text{Me} \\ | \\ \text{O} \\ | \\ \text{N} - \text{C} - \text{CF}_3 \\ | \\ \text{Me} - \text{Si} - \text{CH}_2 - \text{CH} \longrightarrow \text{CH}_2 \\ | \\ \text{Me} \end{array}$$

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Acetamide, N,N-bis(diethylsilyl)-2,2,2-trifluoro- (9CI)
MF C10 H22 F3 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Acetamide, 2,2,2-trifluoro-N,N-bis[tri(methyl-d3)silyl]- (9CI)
MF C8 D18 F3 N O Si2

L12 14 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl ester (9CI)
MF C8 H18 F3 N O Si2
CI COM

$$\begin{array}{c} \text{O-SiMe}_3\\ |\\ \text{Me}_3\text{Si-N---}\text{C-CF}_3 \end{array}$$

ALL ANSWERS HAVE BEEN SCANNED

=> s 112 not (d/els or propenyl or Silanamine )

105525 D/ELS 454965 PROPENYL 6926 SILANAMINE

L13 8 L12 NOT (D/ELS OR PROPENYL OR SILANAMINE )

=> d scan

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Acetamide, N,N-bis(diethylsilyl)-2,2,2-trifluoro- (9CI)
MF C10 H22 F3 N O Si2

$$\begin{array}{c} \text{O} \\ || \\ \text{Et}_2 \text{SiH-N-C-CF}_3 \\ | \\ \text{SiHEt}_2 \end{array}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):10

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Ethanimidic acid, N-(dimethylpropylsily1)-2,2,2-trifluoro-,
dimethylpropylsily1 ester (9CI)
MF C12 H26 F3 N O Si2

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS
IN Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl) - (8CI, 9CI)

MF C8 H18 F3 N O Si2

CI COM

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, N-(diethylsilyl)-2,2,2-trifluoro-, diethylsilyl ester (9CI)

MF C10 H22 F3 N O Si2

$$\begin{array}{c} \text{O-SiHEt}_2 \\ | \\ \text{Et}_2 \text{SiH-N---} \text{C-CF}_3 \end{array}$$

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, N-(ethyldimethylsilyl)-2,2,2-trifluoro-,

ethyldimethylsilyl ester (9CI)

MF C10 H22 F3 N O Si2

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, 2,2,2-trifluoro-N-(triethylsilyl)-, triethylsilyl ester (9CI)

MF C14 H30 F3 N O Si2

$$\begin{array}{c} \text{O-SiEt}_3 \\ | \\ \text{Et}_3 \text{Si-N} \longrightarrow \text{C-CF}_3 \end{array}$$

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, N-[(1,1-dimethylethyl)dimethylsilyl]-2,2,2-trifluoro-,

(1,1-dimethylethyl)dimethylsilyl ester (9CI)

MF C14 H30 F3 N O Si2

L13 8 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl

ester (9CI)

MF C8 H18 F3 N O Si2

CI COM

$$\begin{array}{c} \text{O-SiMe}_3\\ \\ \text{Me}_3\text{Si-N---}\text{C-CF}_3 \end{array}$$

ALL ANSWERS HAVE BEEN SCANNED

```
ANSWER 1 OF 2 REGISTRY COPYRIGHT 2000 ACS
T<sub>1</sub>4
     25561-30-2 REGISTRY
RN
     Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl
CN
     ester (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Acetimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl
     ester (8CI)
     Silanol, trimethyl-, 2,2,2-trifluoro-N-(trimethylsilyl)acetimidate
CN
     (8CI)
OTHER NAMES:
     2,2,2-Trifluoro-N,O-bis(trimethylsilyl)acetamide
     Bis(trimethylsilyl)trifluoroacetamide
CN
CN
     N, O-Bis (trimethylsilyl) trifluoracetamide
CN
CN
     N, O-Bis (trimethylsilyl) trifluoroacetamide
     Trifluoro-N,O-bis(trimethylsilyl)acetamide
CN
     C8 H18 F3 N O Si2
MF
CI
     COM
                 AGRICOLA, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CAPLUS,
LC
     STN Files:
       CASREACT, CHEMCATS, CHEMLIST, CSCHEM, IFICDB, IFIPAT, IFIUDB, MSDS-OHS, PIRA, PROMT, TOXLINE, TOXLIT, USPATFULL
         (*File contains numerically searchable property data)
                       EINECS**
     Other Sources:
          (**Enter CHEMLIST File for up-to-date regulatory information)
           O-SiMe3
Me_3Si-N=C-CF_3
              202 REFERENCES IN FILE CA (1967 TO DATE)
                4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
              202 REFERENCES IN FILE CAPLUS (1967 TO DATE)
     ANSWER 2 OF 2 REGISTRY COPYRIGHT 2000 ACS
L4
     21149-38-2 REGISTRY
RN
     Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI)
CN
     (CA INDEX NAME)
OTHER NAMES:
     N, N-Bis (trimethylsilyl) trifluoroacetamide
CN
FS
     3D CONCORD
     C8 H18 F3 N O Si2
MF
CI
     COM
                   BEILSTEIN*, BIOBUSINESS, CA, CANCERLIT, CAPLUS, CASREACT,
LC
     STN Files:
       CHEMCATS, CHEMLIST, CSCHEM, EMBASE, GMELIN*, HODOC*, IFICDB, IFIPAT,
       IFIUDB, MEDLINE, SPECINFO, TOXLINE, TOXLIT, USPATFULL
         (*File contains numerically searchable property data)
                      EINECS**, NDSL**, TSCA**
     Other Sources:
          (**Enter CHEMLIST File for up-to-date regulatory information)
   MegSi o
Me<sub>3</sub>Si-N-C-CF<sub>3</sub>
```

64 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

more in.

=> dchis\_~

(FILE 'HOME' ENTERED AT 10:07:44 ON 23 JAN 2000)

FILE 'REGISTRY' ENTERED AT 10:07:49 ON 23 JAN 2000

L1 165 S TRIFLUOROACETAMIDE

L2 4 S L1 AND TRIMETHYLSILYL

L3 3 S L2 AND BIS

L4 2 S L3 NOT PMS/CI

FILE 'CAPLUS' ENTERED AT 10:21:13 ON 23 JAN 2000

=> s 14/prep

264 L4

2571797 PREP/RL

L5 8 L4/PREP

(L4 (L) PREP/RL)

=> d ibib abs it hitstr 1-8

L5 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2000 ACS ACCESSION NUMBER: 1999:116666 CAPLUS

ACCESSION NUMBER:

130:153791

DOCUMENT NUMBER: TITLE:

Method for preparation of silylated carboxylic acid

amides

INVENTOR(S):

Winterfeld, Joern; Loskot, Stephan; Menzel, Hartmut;

Reitmeier, Rudolf

PATENT ASSIGNEE(S):

Wacker-Chemie G.m.b.H., Germany

SOURCE:

Ger., 8 pp. CODEN: GWXXAW

(Or

DOCUMENT TYPE: Patent LANGUAGE: German FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE ---------DE 1998-19807021 19980219 DE 19807021 C1 19990211 EP 1998-121577 19981119 EP 937730 19990825 A2 EP 937730 19991013 A3

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

PRIORITY APPLN. INFO.:

DE 1998-19807021 19980219

OTHER SOURCE(S): CASREACT 130:153791

AB The preparation of title compds. via reaction of carboxylic acid chloride with disilazane in the presence of carbonyl group activator is described. Thus, reaction of acetyl chloride with hexamethyldisilazane in the presence of pyridine and Me3SiCl/Me3N gave 90% N,N- as well as N,O-bistrimethylsilylacetamide.

IT Amides, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (silylated; preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane)

TT 78-40-0, Phosphoric acid triethyl ester 80-73-9, 1,3-Dimethyl-2-imidazolidinone 100-71-0, 2-Ethylpyridine 108-48-5, 2,6-Dimethylpyridine 108-99-6 110-86-1, Pyridine, uses 115-86-6, Phosphoric acid triphenyl ester 119-61-9, Benzophenone, uses 126-73-8, Phosphoric acid tributyl ester, uses 603-35-0, Triphenylphosphine, uses 616-47-7, N-Methylimidazole 791-28-6, Triphenylphosphine oxide 872-50-4, N-Methyl-2-pyrrolidinone, uses 998-40-3, Tributylphosphine 19836-78-3, 3-Methyl-2-oxazolidinone 94220-08-3, 1,3-Dimethyltetrahydro-2-pyrimidinone

RL: CAT (Catalyst use); USES (Uses)

(preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane)

TT 75-36-5, Acetyl chloride 79-03-8, Propionic acid chloride 141-75-3, Butyric acid chloride 354-32-5, Trifluoroacetyl chloride 999-97-3, Hexamethyldisilazane 13435-12-6, N-Trimethylsilylacetamide 18156-74-6, N-Trimethylsilylimidazole 18293-54-4 55982-15-5 60498-72-8, N-Trimethylsilyl-2-methylimidazole 119352-64-6 220207-92-1 RL: RCT (Reactant)

(preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane)

IT 10416-58-7P 10416-59-8P 18140-08-4P 21149-38-2P 23728-72-5P 25561-30-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane)

IT 21149-38-2P 25561-30-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane)

RN 21149-38-2 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl) - (8CI, 9CI) (CA INDEX NAME)

Me<sub>3</sub>Si O | || Me<sub>3</sub>Si-N-C-CF<sub>3</sub>

```
ВM
    25561-30-2 CAPLUS
    Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl
CN
    ester (9CI) (CA INDEX NAME)
          O-SiMe3
Me_3Si-N=C-CF_3
    ANSWER 2 OF 8 CAPLUS COPYRIGHT 2000 ACS
                    1998:405958 CAPLUS
ACCESSION NUMBER:
                       129:67865
DOCUMENT NUMBER:
                       Preparation of N,O-bis(trialkylsilyl)perfluoroacylamid
TITLE:
INVENTOR(S):
                       Forat, Gerard
                       Rhodia Chimie, Fr.; Forat, Gerard
PATENT ASSIGNEE(S):
                        PCT Int. Appl., 14 pp.
SOURCE:
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
    PATENT NO.
                    KIND DATE
                                        APPLICATION NO. DATE
                                        -----
     ______
                          -----
                     A1
                          19980618
                                       WO 1997-FR2276 19971211
    WO 9825936
        W: AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GE, HU, ID, IL, IS,
            JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX, NO, NZ, PL, RO,
            SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, AM, AZ, BY, KG, KZ,
            MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI,
            FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM,
            GA, GN, ML, MR, NE, SN, TD, TG
                          19980619
                                        FR 1996-15279
                                                       19961212
     FR 2757167
                     A1
```

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B1
    FR 2757167
                           19990122
                           19980703
                                          AU 1998-54887 19971211
    AU 9854887
                      A1
                          19991006
                                         EP 1997-951313 19971211
    EP 946571
                      A1
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
PRIORITY APPLN. INFO.:
                                          FR 1996-15279
                                                           19961212
                                          WO 1997-FR2276
                                                           19971211
                        CASREACT 129:67865; MARPAT 129:67865
OTHER SOURCE(S):
    The invention concerns the synthesis of N,O-bis(trialkylsilyl)perfluoroacy
     lamides, R1R2R3SiN:C(Rf)OSiR1R2R3 (R1, R2, R3 = C1-10 alkyl; Rf =
     (CX2) pGEA (X = same or different F, CnF2n+1 (n = \leq5), p =
    ≤2; GEA = electron-attracting group, e.g. F or CnF2n+1)) which
    consists in: reacting an amide bearing a Rf group with a trialkylsilyl
    halide in the presence of a base whose halide, or halohydrate, is insol.
     in the medium and in the presence of a nonpolar and nonhydrophilic
     solvent. For example, a mixture of 0.2 mol CF3C(O)NH2 and 0.6 mol Et3N was
     added at 14 mL/h to 0.8 mol Me3SiCl in 120 mL pentane to maintain reflux
     and after addition the mixture was heated at reflux for 11.5 h to yield 89%
    N,O-bis(trimethylsilyl)trifluoroacetamide having 99.5% purity. The
     invention is applicable to organic synthesis.
    Amides, preparation
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
```

(preparation of N,O-bis(trialkylsilyl)perfluoroacylamides)

25561-30-2P, N,O-Bis(trimethylsilyl)trifluoroacetamide

IT

IT

RL: RCT (Reactant)

75-77-4, Chlorotrimethylsilane, reactions 354-38-1, Trifluoroacetamide

(for preparation of N,O-bis(trialkylsilyl)perfluoroacylamides)

```
25561-30-2P, N,O-Bis(trimethylsilyl)trifluoroacetamide
TT
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
    25561-30-2 CAPLUS
RN
    Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl
CN
    ester (9CI) (CA INDEX NAME)
          O-SiMe3
Me_3Si-N=C-CF_3
    ANSWER 3 OF 8 CAPLUS COPYRIGHT 2000 ACS
                        1986:591384 CAPLUS
ACCESSION NUMBER:
                        105:191384
DOCUMENT NUMBER:
                        Simultaneous production of carboxylic acid trimethyl
TITLE:
                        silyl esters and silylated carboxylic acid amides
                        Koetzsch, Hans Joachim; Vahlensieck, Hans Joachim
INVENTOR(S):
                        Dynamit Nobel A.-G., Fed. Rep. Ger.
PATENT ASSIGNEE(S):
                        Ger. Offen., 14 pp.
SOURCE:
                        CODEN: GWXXBX
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        German
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                        APPLICATION NO. DATE
                   KIND DATE
     PATENT NO.
                           -----
                                          ______
     _____ ___
     DE 3443961
                                        DE 1984-3443961 19841201
                           19860605
                     Α1
                     C2
     DE 3443961
                           19861211
                     Α
                                          US 1985-800979
                                                          19851122
                           19860902
     US 4609749
                     A1
                                          GB 1985-29019
                                                          19851125
                           19860604
     GB 2167754
                     B2
                           19890105
     GB 2167754
                                          FR 1985-17465
                                                          19851126
     FR 2574079
                      A1
                           19860606
                     B1
                           19900615
     FR 2574079
                                          JP 1985-267627 19851129
                      A2
                           19860621
     JP 61134393
                                          DE 1984-3443961 19841201
PRIORITY APPLN. INFO.:
     RCO2SiMe3 (I) and RCONHSiMe3 (II) (R = alkyl, alkenyl, haloalkyl), useful
     as silylating agents (no data), are prepared by silylation of (RCO)20 with
     (Me3Si)2NH (III) at 40-130°. Further silylation of II with Me3SiCl
     in the presence of a proton acceptor can give RCON(SiMe3)2 (IV) or
     RC(OSiMe3):NSiMe3 (V). Thus, 660 kg III was treated with 408 kg Ac2O at
     72-90° under N and the resultant mixture was treated with 434 kg
     Me3SiCl in the presence of Me3N at 72° to give 512 kg I (R = Me)
     and a 799 kg mixture of IV and V (R = Me).
IT
     Silylation
        (of anhydrides and amides, with trimethylsilyl chloride and
        hexamethyldisilazane)
                                                         13435-12-6P
     400-53-3P 2754-27-0P
                             10416-58-7P
                                           10416-59-8P
IT
     13688-56-7P 21149-38-2P 25561-30-2P
                                          104890-13-3P
     104890-14-4P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, as silylating agent)
     75-77-4, uses and miscellaneous
IT
     RL: USES (Uses)
        (silylating agent, for amides, silylamide derivs. from)
IT
     999-97-3
     RL: RCT (Reactant)
        (silylating agent, for anhydrides, silylamide derivs. and silyl esters
```

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

from)

IT 108-24-7 407-25-0 760-93-0

RL: RCT (Reactant) (silylation of)

IT 21149-38-2P 25561-30-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as silylating agent)

RN 21149-38-2 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX NAME)

Me<sub>3</sub>Si O | || Me<sub>3</sub>Si N C C CF<sub>3</sub>

RN 25561-30-2 CAPLUS

CN Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl ester (9CI) (CA INDEX NAME)

 $\begin{array}{c} \text{O-SiMe}_3\\ |\\ \text{Me}_3\text{Si-N} \longrightarrow \text{C-CF}_3 \end{array}$ 

L5 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER:

1986:591383 CAPLUS

DOCUMENT NUMBER:

105:191383

TITLE:

Persilylated carboxylic acid amides

INVENTOR(S):

Koetzsch, Hans Joachim; Vahlensieck, Hans Joachim

PATENT ASSIGNEE(S): Dynamit Nobel A.-G., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 11 pp.

\_\_\_\_\_\_

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

			· ·		
	PATENT NO.	KIND	DATE	APPLICATION I	NO. DATE
	DE 3443960	<b>A1</b>	19860605	DE 1984-3443	960 19841201
	DE 3443960	C2	19861211		
	US 4647681	A	19870303	US 1985-8009	96 19851122
	GB 2167753	A1	19860604	GB 1985-2896	0 19851125
	GB 2167753	B2	19890105		
	FR 2574078	A1	19860606	FR 1985-1746	4 19851126
	FR 2574078	B1	19900511		
	JP 61134392	A2	19860621	JP 1985-2662	83 19851128
	JP 08016113	B4	19960221		
PR]	ORITY APPLN. INFO.	:		DE 1984-3443	960 19841201
AB	RCON(SiMe3)2 (I)	and	RC(OSiMe3):NSiMe	3 (TT) (R = H)	. alkvl. alker
MР	KCON (DINC) / 4 (I)	unu	TO (ODTITOD) . HOTING		,

AB RCON(SiMe3)2 (I) and RC(OSiMe3):NSiMe3 (II) (R = H, alkyl, alkenyl, haloalkyl), useful as silylating agents (no data), are prepared by silylation of RCONH2 with 2 equivs. Me3SiCl at 40-80° in the presence of a tertiary amine and (Me3Si)2NH (III). Thus, 9 kg HCONH2, 21.7 kg Me3SiCl, and 140 L III were heated at 48-56° under N in the presence of Me3N to give 92.9% I (R = H).

IT Silylation

(of amides, by trimethylsilyl chloride and hexamethyldilsilazane)

IT 10416-58-7P 15500-60-4P 17898-15-6P 21149-38-2P

26345-88-0P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as silylating agent) 75-77-4, reactions 999-97-3 ΙT RL: RCT (Reactant) (reaction of, with amides, silylamide derivs. from) 60-35-5, reactions 75-12-7, reactions 79-05-0 79-06-1, reactions IT 354-38-1 RL: RCT (Reactant) (silylation of) TT 21149-38-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as silylating agent) RN 21149-38-2 CAPLUS Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX CN Me<sub>3</sub>Si o Me3Si-N-C-CF3 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2000 ACS 1980:106661 CAPLUS ACCESSION NUMBER: 92:106661 DOCUMENT NUMBER: Silylation of tryptamine and 5-hydroxytryptamine: TITLE: kinetic aspects and practical analytical implications Martinez, Emilio; Gelpi, Emilio AUTHOR (S): CORPORATE SOURCE: Inst. Biofis. Neurobiol., CSIC, Barcelona, 25, Spain Adv. Chromatogr. (Houston) (1979), 14th, 677-96 SOURCE: CODEN: ACMGBR DOCUMENT TYPE: Journal English LANGUAGE: An attempt is made to answer some practical kinetic and identification questions on the trimethylsilylated (TMS) derivs. of tryptamine (T) and 5-hydroxytryptamine (5HT). Considering the theor. possible 5 and 11 TMS derivs. of T and 5HT, resp., a gas chromatog.-mass spectrometric study of these reaction profiles established the identification of the 5T derivs. and of 6 of the 11 5HT possibilities. A kinetic approach using different catalysts (trimethylsilylimidazole, trimethylchlorosilane, pyridine) indicated that the ease of silylation would follow the order O5 > 1st  $N\omega$  position > N1 > 2nd  $N\omega$  position. The best results were obtained by reaction with a N,O-bis(trimethylsilyl)trifluoroacetamidetrimethylsilylimidazole-pyridine mixture at 70° fr 60 min, which enhanced the formation of the fully silylated derivs. The relative merits of the silyl vs. the acyl derivs. of these amines are discussed in relation to practical applications. IT Kinetics of silylation (of hydroxytryptamine and tryptamine) IT Mass spectra (of hydroxytryptamine and tryptamine silyl derivs.) IT 75-77-4P, biological studies 343-94-2DP, silyl derivs. 1066-35-9P 10416-59-8P 18156-74-6P **25561-30-2P** 3036-16-6DP, silyl derivs. RL: PREP (Preparation) (preparation of, kinetics of) 25561-30-2P RL: PREP (Preparation) (preparation of, kinetics of) RN25561-30-2 CAPLUS Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl CNester (9CI) (CA INDEX NAME)

```
O-SiMe3
Me_3Si-N=C-CF_3
```

ANSWER 6 OF 8 CAPLUS COPYRIGHT 2000 ACS 1976:524025 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 85:124025 Preparation and structure of trimethylsilyl- and TITLE: trifluoroacetyl-substituted ammonia and hydroxylamine Lidy, Werner; Sundermeyer, Wolfgang AUTHOR (S): Anorg.-Chem. Inst., Univ. Heidelberg, Heidelberg, Ger. CORPORATE SOURCE: Chem. Ber. (1976), 109(7), 2542-6 SOURCE: CODEN: CHBEAM Journal DOCUMENT TYPE: German LANGUAGE: (Me3Si)3N, ClN(SiMe3)2, and Me3SiON(SiMe3)2 reacted with F3CCOCl to give, AB not the expected acylated amines but gave imides, RN:C(CF3)OSiMe3 (R = F3CCO, Cl, Me3SiO, F3CCO2). 60556-43**-**6P 60556-44-7P 25561-30-2P 60556-42-5P IT60556-45-8P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 354-32-5 TΤ RL: RCT (Reactant) (reaction with silylamines) 1070-89-9 1586-73-8 4148-01-0 21023-20-1 TΤ RL: RCT (Reactant) (reaction with trifluoroacetyl chloride) IT25561-30-2P RL: SPN (Synthetic preparation); PREP (Preparation)

 $Me_3Si-N=C-CF_3$ 

ANSWER 7 OF 8 CAPLUS COPYRIGHT 2000 ACS 1970:66309 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

RΝ

CN

Preparation and properties of N-heptafluoroisopropyl and N-trifluoroacetyl sulfur dichloride imide and

sulfur oxide imide, and N,O-

Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl

bis(trimethylsilyl)trifluoroacetimide Von Halasz, Sigmar P.; Glemser, Oskar

CORPORATE SOURCE:

Anorg.-Chem. Inst., Univ. Goettingen, Goettingen, Ger.

Chem. Ber. (1970), 103(2), 553-60 SOURCE:

CODEN: CHBEAM

DOCUMENT TYPE:

AUTHOR(S):

Journal

LANGUAGE: German

(preparation of) 25561-30-2 CAPLUS

ester (9CI) (CA INDEX NAME)

(CF3)2CFN:SC12 was prepared in 54% yield from (CF3)2CFN:SF2 and PC15.(CF3)2C:NH reacted with F2SO to give 23% (CF3)2CFN:SO. CF3CON:SC12 is partly converted by CsF and Cl to CF3CON:SF2. CF3CON:SO was obtained

in 21% yield from Cl2SO and CF3C(OSiMe3):NSiMe3 (I), which was prepared from CF3CONH2 and ClSiMe3 in the presence of NEt3. The 1H- and 19F-NMR spectra suggest the imide structure for I rather than the amide structure. 25561-30-2P 26454-66-0P 26454-67-1P 26454-68-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 25561-30-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 25561-30-2 CAPLUS Ethanimidic acid, 2,2,2-trifluoro-N-(trimethylsilyl)-, trimethylsilyl

O-SiMe3  $Me_3Si-N=C-CF_3$ 

IT

IT

RN

CN

ANSWER 8 OF 8 CAPLUS COPYRIGHT 2000 ACS

ester (9CI) (CA INDEX NAME)

ACCESSION NUMBER:

1969:57999 CAPLUS

DOCUMENT NUMBER:

70:57999

TITLE: INVENTOR(S): Bis(trimethylsilyl)trifluoroacetamide Gehrke, Charles W.; Stalling, David L.

PATENT ASSIGNEE(S):

Research Corp.

SOURCE:

U.S., 1 p. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3415864	A	19681210	US 1967-666975	19670911
GB 1194105	A	19700610	GB 1968-1194105	19680829
PRIORITY APPLN. INFO.			US 1967-666975	19670911
AB Et3N (270 ml.) w	as add	ed to 146 g.	Me3SiCl and the mix	ture slowl

ly added to 58 g. CF3CONH2 under anhydrous conditions and refluxed. The mixture was filtered and distilled in vacuo to yield CF3CON(SiMe3)2 (I), b27 47°. I has application as a reagent for making Me3Si-derivs. of amino acids for gas liquid chromatog.

IT 21149-38-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

21149-38-2P IT

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

21149-38-2 CAPLUS RN

Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX CNNAME)

MegSi o Me<sub>3</sub>Si-N-C-CF<sub>3</sub> L15 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2000 ACS 1992:235863 CAPLUS ACCESSION NUMBER: 116:235863 DOCUMENT NUMBER: Preparation of N,O-bis(tertiary-TITLE: butyldimethylsilyl)trifluroracetamide Endo, Mikio; Ishihara, Toshinobu; Hayashida, Akira; INVENTOR(S): Shinohara, Norio Shin-Etsu Chemical Industry Co., Ltd., Japan PATENT ASSIGNEE(S): Jpn. Kokai Tokkyo Koho, 5 pp. SOURCE: CODEN: JKXXAF Patent DOCUMENT TYPE: Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE KIND DATE PATENT NO. \_\_\_\_\_ ----**---**JP 1990-90687 19900405 19911218 A2 JP 03287592 B2 19980921 JP 2799619 CASREACT 116:235863; MARPAT 116:235863 OTHER SOURCE(S): The title compound (I) was prepared by reaction of CF3CONH2 or CF3CONHSiMe2CMe3 (II) with RMgX (R = hydrocarbyl, X = halo) followed by treatment with Me2CMe2SiCl. Thus, II was treated with MeMgCl in THF at 0-40° for 15 min followed by treatment with Me3CMe2SiCl at 70° for 1.5 h to give 85% I. IT 87020-42-6P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 18162-48-6, tert-Butyldimethylchlorosilane IT RL: RCT (Reactant) (reaction of, with (butyldimethylsilyl)trifluoroacetamide) 83170-48-3 IT 354-38-1 RL: RCT (Reactant) (reaction of, with methylmagnesium chloride and butyldimethylsilyl chloride) 676-58-4, Methylmagnesium chloride IT RL: RCT (Reactant) (reaction of, with trifluoroacetamide derivative and butyldimethylsilyl chloride) IT 87020-42-6P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 87020-42-6 CAPLUS RN

Ethanimidic acid, N-[(1,1-dimethylethyl)dimethylsilyl]-2,2,2-trifluoro-,

(1,1-dimethylethyl)dimethylsilyl ester (9CI) (CA INDEX NAME)

CN

L16 ANSWER 1 OF 3 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

129:67865 CASREACT

TITLE:

Preparation of N,O-bis(trialkylsilyl)perfluoroacylamid

INVENTOR(S):

Forat, Gerard

PATENT ASSIGNEE(S):

Rhodia Chimie, Fr.; Forat, Gerard

SOURCE:

PCT Int. Appl., 14 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE			APPLICATION NO. DATE														
WO S	WO 9825936 A1 19980618				WO 1997-FR2276 19971211												
	W:													ΗU,			
•		JP,	ΚP,	KR,	LC,	LK,	LR,	LT,	LV,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,
		SG,	SI,	SK,	SL,	TR,	TT,	UA,	US,	UΖ,	VN,	YU,	AM,	ΑZ,	BY,	KG,	KZ,
		MD,	RU,	ТJ,	TM												
	RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	ΒE,	CH,	DE,	DK,	ES,	FI,
		FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,
							SN,										
FR 2	2757	167		A	1	1998	0619		F	R 19	96-1	5279		1996	1212		
FR 2	2757	167		В	1	1999	0122										
AU S	98548	387		A.	1	1998	0703		Αl	J 19	98-5	4887		1997	1211		
EP S	9465	71		A	1	1999	1006		E	P 19	97-9	5131	3	1997	1211		
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	FI														
PRIORITY	APP	LN.	INFO	. :					F	R 19	96-1	5279		1996	1212		
									M	0 19	97-F	R227	6	1997	1211		

OTHER SOURCE(S): MARPAT 129:67865

The invention concerns the synthesis of N,O-bis(trialkylsilyl)perfluoroacy lamides, R1R2R3SiN:C(Rf)OSiR1R2R3 (R1, R2, R3 = C1-10 alkyl; Rf = (CX2)pGEA (X = same or different F, CnF2n+1 (n =  $\leq$ 5), p = ≤2; GEA = electron-attracting group, e.g. F or CnF2n+1)) which consists in: reacting an amide bearing a Rf group with a trialkylsilyl halide in the presence of a base whose halide, or halohydrate, is insol. in the medium and in the presence of a nonpolar and nonhydrophilic solvent. For example, a mixture of 0.2 mol CF3C(O)NH2 and 0.6 mol Et3N was added at 14 mL/h to 0.8 mol Me3SiCl in 120 mL pentane to maintain reflux and after addition the mixture was heated at reflux for 11.5 h to yield 89% N,O-bis(trimethylsilyl)trifluoroacetamide having 99.5% purity. The invention is applicable to organic synthesis.

## RX(1) OF 1 2 A В

C YIELD 89%

RCT A 75-77-4, B 354-38-1 RX(1) RGT D 121-44-8 Et3N PRO C **25561-30-2** SOL 109-66-0 Pentane

L16 ANSWER 2 OF 3 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

116:235863 CASREACT

TITLE:

Preparation of N,O-bis(tertiary-

butyldimethylsilyl)trifluroracetamide

INVENTOR(S):

Endo, Mikio; Ishihara, Toshinobu; Hayashida, Akira;

Shinohara, Norio

PATENT ASSIGNEE(S):

Shin-Etsu Chemical Industry Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 03287592	A2	19911218	JP 1990-90687	19900405
TD 2700610	כם	10000021		

JP 2799619

B2 19980921

OTHER SOURCE(S):

MARPAT 116:235863

The title compound (I) was prepared by reaction of CF3CONH2 or CF3CONHSiMe2CMe3 (II) with  $\overline{R}MgX$  (R =  $\overline{h}ydrocarbyl$ , X = halo) followed by treatment with Me2CMe2SiCl. Thus, II was treated with MeMgCl in THF at 0-40° for 15 min followed by treatment with Me3CMe2SiCl at 70° for 1.5 h to give 85% I.

RX(1) OF 2 A + B ===> C

YIELD 85%

RX(1) RCT A 83170-48-3 STAGE(1)

RGT D 676-58-4 MeMgCl

SOL 109-99-9 THF

STAGE (2)

RCT B 18162-48-6 SOL 109-99-9 THF

PRO C 87020-42-6

L16 ANSWER 3 OF 3 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

110:8265 CASREACT

TITLE:

The structure of bis(organosilyl)amides containing the dimethylsilyl and bis(dimethylsilyl)ethylene groups

AUTHOR (S):

Samples, Marjorie S.; Yoder, Claude H.

CORPORATE SOURCE:

Dep. Chem., Franklin and Marshall Coll., Lancaster,

PA, 17604, USA

SOURCE:

J. Organomet. Chem. (1987), 332(1-2), 69-73

CODEN: JORCAI; ISSN: 0022-328X

DOCUMENT TYPE:

LANGUAGE:

Journal English

Bis(dimethylsilyl)amide RCON(SiMe2H)2 (I, R = Et) was prepared by amination AB of Me2SiHCl with EtCONH2 in the presence of Et3N. I (R = Et, Me) and R1CON(SiMe3)2 (II, R1 = H) exist in the amide form and undergo rapid rotation around the C-N bond. II (R1 = Me) exists in the imidate form. The reaction of several bis(trimethylsilyl)amides with Me2SiClCH2CH2SiMe2Cl also produced amides III (R = H, Me, CF3) containing a 5-membered silazane ring. Structures were determined by 13C, 29Si, 14N, and 170 NMR, and free energies of activation for hindered rotation were determined by variable temperature 13C or 29Si NMR. The difference in structure can be attributed to the spatial requirements of the groups attached to Si.

RX(6) OF 7 N + O ===> F...

RCT N 1066-35-9, O 79-05-0 RX(6) RGT P 121-44-8 Et3N PRO F 25561-30-2

SOL 71-43-2 Benzene

## => file beilstein

FILE 'BEILSTEIN' TEMPORARILY UNAVAILABLE SESSION CONTINUES IN FILE 'CASREACT' If this message appears repeatedly, please notify the Help Desk. Enter "HELP STN" for information on contacting the nearest STN Help Desk by telephone or via SEND in the STNMAIL file.

- => s epsilon dielectric constant
  - 539 EPSILON
  - 26 DIELECTRIC
    - 3 DIELECTRICS
  - 29 DIELECTRIC

(DIELECTRIC OR DIELECTRICS)

- 635 CONSTANT
- 81 CONSTANTS
- 698 CONSTANT

(CONSTANT OR CONSTANTS)

L17

O EPSILON DIELECTRIC CONSTANT (EPSILON(W) DIELECTRIC(W) CONSTANT)

- => s dielectric constant
  - 26 DIELECTRIC
  - 3 DIELECTRICS
  - 29 DIELECTRIC

(DIELECTRIC OR DIELECTRICS)

- 635 CONSTANT
- 81 CONSTANTS
- 698 CONSTANT

(CONSTANT OR CONSTANTS)

L18

14 DIELECTRIC CONSTANT

(DIELECTRIC (W) CONSTANT)

=> s 118 and solvent

12249 SOLVENT

4632 SOLVENTS

14789 SOLVENT

(SOLVENT OR SOLVENTS)

L19

10 L18 AND SOLVENT

=> d ibib abs 10

L19 ANSWER 10 OF 10 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

108:75258 CASREACT

TITLE:

Reaction of 2-hydroxydibenzoylmethanes with hydroxylamine hydrochloride in DMF-water.

Dielectric constant dependent

products: isoxazole and benzisoxazole

AUTHOR(S):

Naphade, V. J.; Ghiya, B. J.

CORPORATE SOURCE:

Inst. Sci., Nagpur, 440 001, India

SOURCE:

J. Indian Chem. Soc. (1986), 63(9), 851-3

CODEN: JICSAH; ISSN: 0019-4522

DOCUMENT TYPE:

LANGUAGE:

Journal

GΙ

English

AB Hydroxydibenzoylmethanes (I; R1, R1 given: Me, MeO; Me, H; H, MeO; H, H) react with NH2OH·HCl in DMF, aqueous DMF and N-methylacetamide to yield isoxazoles and benzisoxazoles. Solvent effects on product yields were discussed.

=> d ibib abs 9

L19 ANSWER 9 OF 10 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

118:21809 CASREACT

TITLE:

Kinetics of oxidation of thioacetic acid by methylene

blue in methanol-water medium

AUTHOR (S):

Chansoria, K.; Mishra, K. K.

CORPORATE SOURCE:

Dep. Postgrad. Stud. Res. Chem., R D Univ., Jabalpur,

482 001, India

SOURCE:

Indian J. Chem., Sect. A: Inorg., Bio-inorg., Phys.,

Theor. Anal. Chem. (1992), 31A(9), 728-31

CODEN: ICACEC

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The title reaction kinetics, to give the coresponding disulfide and the dye leuco base were determined The pseudo 2-nd order rate constant increases linearly with the H+ concentration; the 0.5 rate constant has a fractional order

dependence on the H+ concentration at low H+ concns. is again linear in H+ concentration

at  $\geq 4.0 \times 10^{-2}$  mol dm-3. The pseudo 2-nd order rate constant decreases to a limiting value as the mediums dielec. constant decreases; the 0.5 order process continuously decreases. Me methacrylate does not affect the initial reaction rate; it retards rate of the latter part of the reaction.

=> d ibib abs 6-8

L19 ANSWER 6 OF 10 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

119:249389 CASREACT

TITLE:

Medium effect on the kinetics and mechanism of oxidation of thiocarbohydrazide by chloramine-B Gowda, B. Thimme; Panicker, B. K.; Pardhasaradhi, V.

AUTHOR(S): CORPORATE SOURCE:

Dep. Post-Grad. Stud. Res. Chem., Mangalore Univ.,

Mangalagangothri, 574 199, India

SOURCE: Oxid. Commun. (1993), 16(1-2), 44-61

CODEN: OXCODW; ISSN: 0209-4541

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Medium effect on the kinetics and mechanism of oxidation of thiocarbohydrazide (TCH) by chloramine-B has been investigated in various binary solvent mixts. of varying compns. ranging from 1:9 to 7:3. The solvent mixts. employed are aquo-methanol, aquo-i-propanol and aquo-t-butanol. The rate dependences in

[oxidant], [TCH] and [H+] have been determined in each of these media under varying compns. The rates have also been determined at varying temperature in

all

the compns. Mechanisms consistent with the observed results are discussed. The coeffs. of the rate determining steps have been calculated as functions of solvent, their composition and temperature. The activation parameters corresponding to these consts. are also calculated in all the solvents at varying solvent compns. Both the rate coeffs. and the activation parameters under varying conditions have been correlated. The validity of isokinetic relationship has also been tested.

L19 ANSWER 7 OF 10 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

119:202916 CASREACT

TITLE:

Proton-transfer dynamics in substituted 3-hydroxyflavones: **solvent** polarization

effects

AUTHOR (S):

Swinney, T. C.; Kelley, D. F.

CORPORATE SOURCE:

Dep. Chem., Colorado State Univ., Fort Collins, CO,

80523, USA

SOURCE:

rates

J. Chem. Phys. (1993), 99(1), 211-21

CODEN: JCPSA6; ISSN: 0021-9606

DOCUMENT TYPE:

Journal

LANGUAGE:

TAGE: English

The spectroscopy and excited state proton transfer (ESPT) dynamics of

4'-N,N-dimethylamino-3-hydroxyflavone and 4'-N,N-diethylamino-3-hydroxyflavone have been studied in acetonitrile/benzene solvent mixts. Solvent composition-dependent spectral shifts are observed and can be understood in terms of an Onsager cavity model. Anal. of these spectral shifts accurately predicts solvent composition-dependent excited state equilibrium consts., which are also exptl. determined The ESPT

are analyzed within the framework of a transition state theory treatment of **solvent** polarization-mediated proton transfer. This treatment is analogous to electron transfer theory. In this treatment, the energetics of the transition state are largely determined by known **solvent** properties and the **solvent**-dependent spectroscopy. This anal. yields **solvent**-dependent ESPT activation energies. The corresponding calculated ESPT rates are in excellent agreement with the exptl. determined rates.

L19 ANSWER 8 OF 10 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER:

118:38252 CASREACT

TITLE:

Biomimetic catalysis of SN2 reactions through cation- $\pi$  interactions. The role of polarizability in catalysis

AUTHOR(S):

McCurdy, Alison; Jimenez, Leslie; Stauffer, David A.;

Dougherty, Dennis A.

CORPORATE SOURCE:

Arnold and Mabel Beckman Lab. Chem. Synth., California

Inst. Technol., Pasadena, CA, 91125, USA
J. Am. Chem. Soc. (1992), 114(26), 10314-21

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

Journal

LANGUAGE:

SOURCE:

English

GΙ

$$Cs + O_2C$$
 $CO_2 - Cs^+$ 
 $Cs + O_2C$ 
 $CO_2 - Cs^+$ 
 $Cs + O_2C$ 
 $Co_2 - Cs^+$ 
 $CH_2$ 
 $Cs + O_2C$ 
 $Co_2 - Cs^+$ 
 $CH_2$ 
 $CH_2$ 
 $CH_2$ 

Cyclophane hosts I and II are effective catalysts for both the alkylation of quinoline structures to produce quinolinium salts and the dealkylation of sulfonium salts to produce sulfides. Reactions that develop pos. charge in the transition state and reactions that destroy pos. charge are accelerated by I or II. The former observation is not surprising, given the well-documented ability of these hosts to bind cations through the cation- $\pi$  interaction. The catalysis of the dealkylation reactions, however, along with several other observations, suggests that some other factor is involved in the catalysis. It is proposed that the high polarizability of the transition states is well matched to the very polarizable hosts and that this contributes to the catalysis.

=> s pnictine

1 PNICTINE

=> d ibib abs

L20 ANSWER 1 OF 1 CASREACT COPYRIGHT 2000 ACS

ACCESSION NUMBER: 119:95818 CASREACT

TITLE: Preparation of organosilanes from catalytic silylation

of organic halides with disilanes

INVENTOR(S): Babin, Pierre; Bennetau, Bernard; Dunogues, Jacques

PATENT ASSIGNEE(S): Rhone-Poulenc Chimie, Fr.

SOURCE: Fr. Demande, 14 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	<b>-</b>			
FR 2677358	A1	19921211	FR 1991-6787	19910605
FR 2677358	B1	19940930		

OTHER SOURCE(S): MARPAT 119:95818

AB Organosilyl derivs. are prepared by reaction of ArX [Ar = certain carbocycles (un)substituted by F, -CN, perfluoroalkyl, or Ar = certain

nitrogen heterocycles; X = Cl, I, or preferably Br] with disilanes R1R2R3SiSiR4R5R6 [R1-R6 = hydrocarbon chain, (un)functionalized cycloalkyl or aralkyl, Cl, with the condition that at least one of R1 and R4 ≠ Cl] in the presence of a catalyst comprising a Group VIII metal prictine complex (platinum-group metal with prictine being a phosphine preferred) with or without solvent (preferably an aprotic solvent, e.g., hexamethylphosphorotriamide). Thus, reaction of 10 mmol 4-F3CC6H4Br with 17 mmol (EtO)2SiMeSiMe(OEt)2 for 15 h at 160° in a dried tube in the presence of palladium catalyst (unspecified) gave 4-F3CC6H4SiMe(OEt)2 in 70% yield. A number of trimethylsilyl-substituted benzene and heterocycle derivs. were prepared by this method in high yields.

```
=> s 14/prep
```

175 L4

2642631 PREP/RL

10 L4/PREP L5

(L4 (L) PREP/RL)

=> d ibib abs hitstr 1-10

ANSWER 1 OF 10 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER:

1999:116666 CAPLUS

DOCUMENT NUMBER:

130:153791

TITLE:

Method for preparation of silylated carboxylic acid

INVENTOR(S):

Winterfeld, Joern; Loskot, Stephan; Menzel, Hartmut;

Reitmeier, Rudolf

PATENT ASSIGNEE(S):

Wacker-Chemie G.m.b.H., Germany

SOURCE:

Ger., 8 pp. CODEN: GWXXAW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT	NO.		KII	ИD	DATE			AP	PL:	CATI	ои ис	ο.	DATE			
DE	1980	7021		C:	1	1999	0211		DE	19	998-1	9807	021	1998	0219		
EΡ	9377	730	*	A2	2	1999	0825		EP	19	998-1	2157	7	1998	1119		
$\mathbf{EP}$	9377	730		A.	3	1999	1013								•		
	-	200	DE	CITT	D.17	DIZ	D.C.	TO D	CD	CD	TIT	тт	TTT	TTA	CE	MC	1

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

DE 1998-19807021 19980219 PRIORITY APPLN. INFO.:

CASREACT 130:153791 OTHER SOURCE(S): The preparation of title compds. via reaction of carboxylic acid chloride with disilazane in the presence of carbonyl group activator is described. Thus, reaction of acetyl chloride with hexamethyldisilazane in the presence of pyridine and Me3SiCl/Me3N gave 90% N,N- as well as N, O-bistrimethylsilylacetamide.

IT21149-38-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of silylated carboxylic acid amide via carbonyl group activator promoted reaction of carboxylic acid chloride with disilazane) 21149-38-2 CAPLUS

Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX CN NAME)

Me<sub>3</sub>Si o Me<sub>3</sub>Si-N-C-CF<sub>3</sub>

ANSWER 2 OF 10 CAPLUS COPYRIGHT 2000 ACS ACCESSION NUMBER:

1993:60312 CAPLUS

DOCUMENT NUMBER:

118:60312

Manufacture of heat- and chemically resistant imides TITLE:

INVENTOR(S):

Nozaki, Choji Fuji Photo Film Co., Ltd., Japan

PATENT ASSIGNEE(S): SOURCE:

Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE JP 04187673 A2 19920706 JP 1990-319746 19901122

The title imides (poly-, bis-, and monoimides) are manufactured by treating amic acids with trialkylsilylating agents. Thus, a AcNMe2 solution of 0.137 g biphenyltetracarboxylic acid-bisphenol AF polyamic acid and 1.0 g  $\,$ CF3CONMeSiMe3 was coated on a KBr plate and heated in vacuo at 100° for 1 h to give a silylated polyimide.

24589-78-4DP, N-Methyl-N-(trimethylsilyl)trifluoroacetamide, IT reaction products with amic acids

RL: PREP (Preparation)

(chemical and heat-resistant, manufacture of)

24589-78-4 CAPLUS RN

Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) (CA INDEX NAME)

MegSi o Me-N-C-CF3

ANSWER 3 OF 10 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER: 1990:405668 CAPLUS

DOCUMENT NUMBER: TITLE:

On some new trifluoromethyl iodine(III) compounds:

reactions of CF3IF2 with boron and silicon compounds

and CF3ICl2 with silver salts Tyrra, Wieland; Naumann, Dieter

AUTHOR (S): CORPORATE SOURCE:

Fachbereich Chem., Univ. Dortmund, Dortmund,

D-4600/50, Fed. Rep. Ger.

SOURCE:

J. Fluorine Chem. (1989), 45(3), 401-16

CODEN: JFLCAR; ISSN: 0022-1139

DOCUMENT TYPE:

Journal

LANGUAGE:

English

113:5668

OTHER SOURCE(S):

CASREACT 113:5668

CF3IF2 undergoes fluorine exchange reactions with BX3 (X = Cl, Br, I, OCOCF3) to form CF3IX2. The reactions of CF3IF2 with (CF3)2BN(CH3)2, (CH3)3SiNCO and (CH3)3SiN(CH3)COCF3 yield the corresponding new trifluoromethyl iodine(III) nitrogen compds. CF3ICl2 was prepared by reacting CF3IF2 with (CH3)3SiCl. CF3ICl2 reacts with AgX (X = O2CCF3, SCF3) to yield the corresponding CF3IX2 compds.; with (C6H5)4AsCl, the novel ion [CF3ICl3] - is detected. Products were identified by NMR. CF3I(NCO)2 explosively decomps. when warmed to room temperature

IT 24589-78-4P

RL: FORM (Formation, nonpreparative); PREP (Preparation) (formation of, in reactions of difluoro(trifluomethyl)iodine)

24589-78-4 CAPLUS RN

Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) (CA CNINDEX NAME)

Me<sub>3</sub>Si o Me-N-C-CF3 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2000 ACS

1987:496883 CAPLUS ACCESSION NUMBER:

107:96883 DOCUMENT NUMBER:

N-Methyl-N-(trimethylsilyl)trifluoroacetamide TITLE:

Shinohara, Norio; Inoue, Yoshibumi INVENTOR(S):

Shin-Etsu Chemical Industry Co., Ltd., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 3 pp. SOURCE:

CODEN: JKXXAF

Patent DOCUMENT TYPE:

Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 62042991	A2	19870224	JP 1985-182740	19850820
JP 63027354	B4	19880602		
US 32717	E	19880719	US 1987-109775	19871019
TORITY APPLN. INFO.:	:		JP 1985-182740	19850820
			US 1986-896280	19860814

The title compound (I), useful as a reagent for antibiotics synthesis (no AB data), is prepared by heating F3CCONHMe (II) with N,Obis(trimethylsilyl)acetamide (III). A solution of 254 g II and 258 g III in 400 mL C6H6 was stirred at 80° for 3 h to give 91% I.

24589-78-4P, N-Methyl-N-(trimethylsilyl)trifluoroacetamide TT

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of, as silylating agent for antibiotic synthesis)

RN 24589-78-4 CAPLUS

Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) CN INDEX NAME)

PR

ANSWER 5 OF 10 CAPLUS COPYRIGHT 2000 ACS

1987:459249 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 107:59249

Method for the preparation of N-methyl-N-TITLE:

(trimethylsilyl)trifluoroacetamide

Shinohara, Toshio; Inoue, Yoshifumi INVENTOR(S):

Shin-Etsu Chemical Industry Co., Ltd., Japan PATENT ASSIGNEE(S):

U.S., 3 pp. SOURCE:

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				<b></b>
US 4663471	A	19870505	US 1986-896280	19860814
US 32717	E	19880719	US 1987-109775	19871019
PRIORITY APPLN. INF	0.:	•	JP 1985-182740	19850820
			US 1986-896280	19860814

The title compound (I) is prepared by silylation of N-methyltrifluoroacetamide AΒ (II) with N,O-bis(trimethylsilyl)acetamide (III) at 40-130°. I was produced in 91% yield by reaction of 2 mol II and 2.6 mol III at

80° for 3 h. 24589-78-4P IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by silylation of methyltrifluoroacetamide with N,O-bis(trimethylsilyl)acetamide)

RN24589-78-4 CAPLUS

Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) (CA CNINDEX NAME)

Me<sub>3</sub>Si o Me-N-C-CF3

ANSWER 6 OF 10 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER:

1986:591384 CAPLUS

DOCUMENT NUMBER:

105:191384

TITLE:

Simultaneous production of carboxylic acid trimethyl silyl esters and silylated carboxylic acid amides Koetzsch, Hans Joachim; Vahlensieck, Hans Joachim

INVENTOR(S):

Dynamit Nobel A.-G., Fed. Rep. Ger.

PATENT ASSIGNEE(S):

Ger. Offen., 14 pp.

SOURCE:

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	API	LICATION NO.	DATE
DE 3443961	<b>A</b> 1	19860605	DE	1984-3443961	19841201
DE 3443961	C2	19861211			
US 4609749	Α	19860902	US	1985-800979	19851122
GB 2167754	A1	19860604	GB	1985-29019	19851125
GB 2167754	B2	19890105			
FR 2574079	A1	19860606	FR	1985-17465	19851126
FR 2574079	B1	19900615			
JP 61134393	A2	19860621	JΡ	1985-267627	19851129
PRIORITY APPLN. INFO.:			DE	1984-3443961	19841201

RCO2SiMe3 (I) and RCONHSiMe3 (II) (R = alkyl, alkenyl, haloalkyl), useful as silylating agents (no data), are prepared by silylation of (RCO)20 with (Me3Si)2NH (III) at 40-130°. Further silylation of II with Me3SiCl in the presence of a proton acceptor can give RCON(SiMe3)2 (IV) or RC(OSiMe3):NSiMe3 (V). Thus, 660 kg III was treated with 408 kg Ac2O at 72-90° under N and the resultant mixture was treated with 434 kg Me3SiCl in the presence of Me3N at 72° to give 512 kg I (R = Me) and a 799 kg mixture of IV and V (R = Me).

21149-38-2P IT

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as silylating agent)

21149-38-2 CAPLUS RN

Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX CN NAME)

L5 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER: 1986:591383 CAPLUS

DOCUMENT NUMBER:

105:191383

TITLE:

Persilylated carboxylic acid amides

INVENTOR (S):

Koetzsch, Hans Joachim; Vahlensieck, Hans Joachim

PATENT ASSIGNEE(S):

Dynamit Nobel A.-G., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 11 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	<b>-</b>			
DE 3443960	A1	19860605	DE 1984-3443960	19841201
DE 3443960	C2	19861211		
US 4647681	Α	19870303	US 1985-800996	19851122
GB 2167753	A1	19860604	GB 1985-28960	19851125
GB 2167753	B2	19890105		
FR 2574078	A1	19860606	FR 1985-17464	19851126
FR 2574078	B1	19900511		
JP 61134392	A2	19860621	JP 1985-266283	19851128
JP 08016113	B4	19960221		
			DD 1004 2442060	10041001

PRIORITY APPLN. INFO .:

DE 1984-3443960 19841201

AB RCON(SiMe3)2 (I) and RC(OSiMe3):NSiMe3 (II) (R = H, alkyl, alkenyl, haloalkyl), useful as silylating agents (no data), are prepared by silylation of RCONH2 with 2 equivs. Me3SiCl at 40-80° in the presence of a tertiary amine and (Me3Si)2NH (III). Thus, 9 kg HCONH2, 21.7 kg Me3SiCl, and 140 L III were heated at 48-56° under N in the presence of Me3N to give 92.9% I (R = H).

IT 21149-38-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of, as silylating agent)

RN 21149-38-2 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX NAME)

Me<sub>3</sub>Si O | || Me<sub>3</sub>Si-N-C-CF<sub>3</sub>

L5 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2000 ACS

ACCESSION NUMBER:

1977:467465 CAPLUS

DOCUMENT NUMBER:

87:67465

TITLE:

Orientation effects in charge transfer interactions.

11. Stereoisomeric quinhydrones of the

[2.2] (1,4) naphthalenophane series

AUTHOR(S):

Staab, Heinz A.; Herz, Claus P.

CORPORATE SOURCE:

Abt. Org. Chem., Max-Planck-Inst. Med. Forsch.,

Heidelberg, Ger.

SOURCE:

Angew. Chem. (1977), 89(6), 406-7

CODEN: ANCEAD

DOCUMENT TYPE:

Journal

LANGUAGE:

German

GI For diagram(s), see printed CA Issue.

AB Both I and II show similar charge-transfer bands in the 500-700 nm region; these results are rationalized in terms of a weak donor-acceptor interaction through space in these systems.

IT 24589-78-4P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 24589-78-4 CAPLUS Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) RNCNINDEX NAME) Me<sub>3</sub>Si o Me-N-C-CF3 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2000 ACS 1972:59694 CAPLUS ACCESSION NUMBER: 76:59694 DOCUMENT NUMBER: Organosilicon compounds. XLVI. New synthesis of TITLE: heptamethyldisilazane and N-methyl-N-(trimethylsilyl)trifluoroacetamide

heptamethyldisilazane and N-methyl-N(trimethylsilyl)trifluoroacetamide

AUTHOR(S): Birkofer, Leonhard; Schmidtberg, Guenter

CORPORATE SOURCE: Inst. Org. Chem., Univ. Duesseldorf, Duesseldorf, Ger.

SOURCE: Chem. Ber. (1971), 104(12), 3831-4

CODEN: CHBEAM

DOCUMENT TYPE: Journal LANGUAGE: German

AB Me3SiCl and MeNH2 in pentane gave 85% (Me3Si)2NMe, which reacted with (F3CCO)2O at room temperature to give 95% F3CCON(Me)SiMe3.

IT 24589-78-4P
RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 24589-78-4 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N-methyl-N-(trimethylsilyl)- (8CI, 9CI) (CA INDEX NAME)

Me<sub>3</sub>Si O | || Me-N-C-CF<sub>3</sub>

L5 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2000 ACS ACCESSION NUMBER: 1969:57999 CAPLUS

DOCUMENT NUMBER: 70:57999

TITLE: Bis(trimethylsilyl)trifluoroacetamide
INVENTOR(S): Gehrke, Charles W.; Stalling, David L.

PATENT ASSIGNEE(S): Research Corp. SOURCE: U.S., 1 p.

CODEN: USXXAM
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 3415864 A 19681210 US 1967-666975 19670911
GB 1194105 A 19700610 GB 1968-1194105 19680829
PRIORITY APPLN. INFO.: US 1967-666975 19670911

AB Et3N (270 ml.) was added to 146 g. Me3SiCl and the mixture slowly added to 58 g. CF3CONH2 under anhydrous conditions and refluxed. The mixture was

filtered and distilled in vacuo to yield CF3CON(SiMe3)2 (I), b27 47°. I has application as a reagent for making Me3Si-derivs. of amino acids for gas liquid chromatog.

IT 21149-38-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 21149-38-2 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N,N-bis(trimethylsilyl)- (8CI, 9CI) (CA INDEX NAME)

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=> s pentafluoropropanoic acid/cn

1 PENTAFLUOROPROPANOIC ACID/CN

=> d scan

L6 1 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Propanoic acid, pentafluoro- (9CI)

MF C3 H F5 O2

CI COM

 $HO_2C-CF_2-CF_3$ 

=> s c3h2f5no/mf

L7 2 C3H2F5NO/MF

=> d scan

L7 2 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Propanamide, 2,2,3,3,3-pentafluoro- (9CI)

MF C3 H2 F5 N O

CI COM

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L7 2 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Propanimidoyl fluoride, 2,3,3,3-tetrafluoro-N-hydroxy- (9CI)

MF C3 H2 F5 N O

CI COM

## ALL ANSWERS HAVE BEEN SCANNED

=> s propanamide

L8 80775 PROPANAMIDE

=> s propanamide and 17

80775 PROPANAMIDE

L9 1 PROPANAMIDE AND L7

 $\Rightarrow$  s 18 and 1-3/si

849289 1-3/SI

L10 879 L8 AND 1-3/SI

=> s 18 and 1-2/si

819660 1-2/SI

L11 855 L8 AND 1-2/SI

=> s lll and 1/nc

22233131 1/NC

L12 817 L11 AND 1/NC

=> d scan

L12 817 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN 1-Piperidinepropanamide, N-[4-[[(1,1-dimethylethyl)dimethylsilyl]oxy]
 phenyl]-4-hydroxy-4-(phenylmethyl)- (9CI)

MF C27 H40 N2 O3 Si

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 18 and 1/n

3338029 1/N L13 17260 L8 AND 1/N

=> s 112 and 1/n

3338029 1/N

L14 383 L12 AND 1/N

=> s l14 and f/els

1422047 F/ELS

L15 30 L14 AND F/ELS

=> d scan

L15 30 ANSWERS REGISTRY COPYRIGHT 2000 ACS

IN Propanamide, 2,3,3,3-tetrafluoro-2-[1,1,2,3,3,3-hexafluoro-2-(heptafluoropropoxy)propoxy]-N-[3-(trimethoxysily1)propy1]- (9CI)

MF C15 H16 F17 N O6 Si

CI COM